

## CLAIMS

### What is Claimed Is:

Claim 1. A method for treating a patient suffering from a cancerous disease comprising:

administering to said patient an anti-cancer antibody or fragment thereof produced in accordance with a method for the production of anti-cancer antibodies which are useful in treating a cancerous disease, said antibody or fragment thereof characterized as being cytotoxic against cells of a cancerous tissue, and being essentially benign to non-cancerous cells;

wherein said antibody or fragment thereof is placed in admixture with a pharmaceutically acceptable adjuvant and is administered in an amount effective to mediate treatment of said cancerous disease;

said antibody being an isolated monoclonal antibody or antigen binding fragment thereof which binds to an antigenic moiety expressed by said cancerous tissue, said antigenic moiety characterized as being bound by an antibody having identifying characteristics of a monoclonal antibody encoded by a clone deposited with the ATCC as PTA-4890.

Claim 2. The method for treating a patient suffering from a cancerous disease in accordance with claim 1, wherein said antibody or fragment thereof is humanized or chimerized.

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2           Claim 3.       The method for treating a patient suffering from a cancerous disease  
3 in accordance with claim 1 comprising:

4           conjugating said antibody or antigen binding fragment thereof with a member  
5 selected from the group consisting of toxins, enzymes, radioactive compounds, and  
6 hematogenous cells, thereby forming an antibody conjugate; and

7           administering said antibody conjugate or conjugated fragments thereof to said  
8 patient;

9           wherein said antibody conjugate or conjugated fragments are placed in admixture  
10 with a pharmaceutically acceptable adjuvant and are administered in an amount effective to  
11 mediate treatment of said cancerous disease.

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13           Claim 4.       The method of claim 3, wherein said antibody or fragment thereof is  
14 humanized or chimerized.

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1           Claim 5. The method for treating a patient suffering from a cancerous disease in  
2 accordance with claim 1 wherein:

3           the cytotoxicity of said antibody or fragment thereof is mediated through  
4 antibody dependent cellular toxicity.

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6           Claim 6. The method for treating a patient suffering from a cancerous disease in  
7 accordance with claim 1 wherein:

8           the cytotoxicity of said antibody or fragment thereof is mediated through  
9 complement dependent cellular toxicity.

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11           Claim 7. The method for treating a patient suffering from a cancerous disease in  
12 accordance with claim 1 wherein:

13           the cytotoxicity of said antibody or fragment thereof is mediated through  
14 catalyzing of the hydrolysis of cellular chemical bonds.

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16           Claim 8. The method for treating a patient suffering from a cancerous disease in  
17 accordance with claim 1 wherein:

18           the cytotoxicity of said antibody or fragment thereof is mediated through  
19 producing an immune response against putative cancer antigens residing on tumor  
20 cells.

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1           Claim 9. The method for treating a patient suffering from a cancerous disease in  
2   accordance with claim 1 wherein:

3           the cytotoxicity of said antibody or fragment thereof is mediated through  
4   targeting of cell membrane proteins to interfere with their function.

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6           Claim 10. The method for treating a patient suffering from a cancerous disease  
7   in accordance with claim 1 wherein:

8           the cytotoxicity of said antibody or fragment thereof is mediated through  
9   production of a conformational change in a cellular protein effective to produce a  
10   signal to initiate cell-killing.

11

12           Claim 11.     The method for treating a patient suffering from a cancerous  
13   disease in accordance with claim 1 wherein:

14           said method of production utilizes a tissue sample containing cancerous and  
15   non-cancerous cells obtained from a particular individual.

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17           Claim 12.     A method for treating a patient suffering from a cancerous  
18   disease comprising:

19           administering to said patient an antibody or antigen binding fragment thereof  
20   produced in accordance with a method for the production of anti-cancer antibodies  
21   which are useful in treating a cancerous disease, said antibody being cytotoxic against  
22   cells of a cancerous tissue, and essentially benign to non-cancerous cells;

1            wherein said antibody is the isolated monoclonal antibody encoded by the clone  
2       deposited with the ATCC as PTA-4890 or an antigen binding fragment thereof, and is  
3       placed in admixture with a pharmaceutically acceptable adjuvant and is administered in  
4       an amount effective to mediate treatment of said cancerous disease.

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6            Claim 13.     The method for treating a patient suffering from a cancerous  
7       disease in accordance with claim 12, wherein said antibody or fragment thereof is  
8       humanized or chimerized.

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10           Claim 14.     The method for treating a patient suffering from a cancerous  
11       disease in accordance with claim 12 comprising:

12           conjugating said antibody or fragment thereof with a member selected from the  
13       group consisting of toxins, enzymes, radioactive compounds, and hematogenous cells,  
14       whereby an antibody conjugate is formed; and

15           administering said antibody conjugates or fragments thereof to said patient;

16           wherein said conjugated antibodies are placed in admixture with a  
17       pharmaceutically acceptable adjuvant and are administered in an amount effective to  
18       mediate treatment of said cancerous disease.

19           Claim 15.     The method of claim 14, wherein said antibody or fragment  
20       thereof is selected from said subset are humanized or chimerized.

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1           Claim 16. The method for treating a patient suffering from a cancerous disease  
2   in accordance with claim 12 wherein:

3           the cytotoxicity of said antibody or fragment thereof is mediated through  
4   antibody dependent cellular toxicity.

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6           Claim 17. The method for treating a patient suffering from a cancerous disease  
7   in accordance with claim 12 wherein:

8           the cytotoxicity of said antibody or fragment thereof is mediated through  
9   complement dependent cellular toxicity.

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11          Claim 18. The method for treating a patient suffering from a cancerous disease  
12   in accordance with claim 12 wherein:

13          the cytotoxicity of said antibody or fragment thereof is mediated through  
14   catalyzing of the hydrolysis of cellular chemical bonds.

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16          Claim 19. The method for treating a patient suffering from a cancerous disease  
17   in accordance with claim 12 wherein:

18          the cytotoxicity of said antibody or fragment thereof is mediated through  
19   producing an immune response against putative cancer antigens residing on tumor  
20   cells.

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1           Claim 20. The method for treating a patient suffering from a cancerous disease  
2   in accordance with claim 12 wherein:

3           the cytotoxicity of said antibody or fragment thereof is mediated through  
4   targeting of cell membrane proteins to interfere with their function.

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6           Claim 21. The method for treating a patient suffering from a cancerous disease  
7   in accordance with claim 12 wherein:

8           the cytotoxicity of said antibody or fragment thereof is mediated through  
9   production of a conformational change in a cellular protein effective to produce a  
10   signal to initiate cell-killing.

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12           Claim 22.     The method for treating a patient suffering from a cancerous  
13   disease in accordance with claim 12 wherein:

14           said method of production utilizes a tissue sample containing cancerous and  
15   non-cancerous cells obtained from a particular individual.

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17           Claim 23.     A process for mediating cytotoxicity of a human tumor cell  
18   which expresses a CD63 antigenic moiety on the cell surface comprising:

19                   contacting said tumor cell with an isolated monoclonal antibody or  
20   antigen binding fragment thereof, said antibody or antigen binding fragment thereof  
21   being an isolated monoclonal antibody or antigen binding fragment thereof which  
22   binds to said expressed CD63 antigenic moiety, said antigenic moiety characterized as

1 being bound by an antibody having the identifying characteristics of a monoclonal  
2 antibody encoded by the clone deposited with the ATCC as PTA-4890,  
3 whereby cell cytotoxicity occurs as a result of said binding.

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5 Claim 24. The process of claim 23 wherein said isolated antibody or  
6 antigen binding fragments thereof are humanized or chimerized.

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8 Claim 25. The process of claim 23 wherein said isolated antibody or  
9 antigen binding fragments thereof are conjugated with a member selected from the  
10 group consisting of cytotoxic moieties, enzymes, radioactive compounds, and  
11 hematogenous cells, whereby an antibody conjugate is formed..

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13 Claim 26. The process of claim 23 wherein said isolated antibody or  
14 antigen binding fragments thereof are humanized or chimerized.

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16 Claim 27. The process of claim 23 wherein said isolated antibody or  
17 antigen binding fragments thereof are murine.

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19 Claim 28. The process of claim 23 wherein the human tumor tissue sample  
20 is obtained from a tumor originating in a tissue selected from the group consisting of  
21 colon, ovarian, lung, prostate and breast tissue.

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1           Claim 29.     A binding assay to determine a presence of cells which express a  
2     CD63 antigenic moiety which specifically binds to an isolated monoclonal antibody  
3     encoded by the clone deposited with the ATCC as PTA-4890 or an antigen binding  
4     fragment thereof comprising:

5           providing a cell sample;

6           providing an isolated monoclonal antibody or antigen binding fragment thereof,  
7     said antibody or antigen binding fragment thereof being an isolated monoclonal  
8     antibody or antigen binding fragment thereof which binds to said expressed CD63  
9     antigenic moiety, said antigenic moiety characterized as being bound by an antibody  
10    having the identifying characteristics of a monoclonal antibody encoded by the clone  
11    deposited with the ATCC as PTA-4890;

12          contacting said isolated monoclonal antibody or antigen binding fragment  
13    thereof with said cell sample; and

14          determining binding of said isolated monoclonal antibody or antigen binding  
15    fragment thereof with said cell sample;

16          whereby the presence of cells which express a CD63 antigenic moiety which  
17    specifically binds to said isolated monoclonal antibody or antigen binding fragment  
18    thereof is determined.

19          Claim 30.     The binding assay of claim 29 wherein the cell sample is  
20    obtained from a tumor originating in a tissue selected from the group consisting of  
21    colon, ovarian, lung, prostate and breast tissue.

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1           Claim 31.     A process of isolating or screening for cells in a sample which  
2   express a CD63 antigenic moiety which specifically binds to an isolated monoclonal  
3   antibody or antigen binding fragment thereof, said antigenic moiety characterized as  
4   being bound by an antibody having the identifying characteristics of a monoclonal  
5   antibody encoded by the clone deposited with the ATCC as PTA-4890 comprising:  
  
6           providing a cell sample;  
  
7           providing an isolated monoclonal antibody or antigen binding fragment thereof,  
8   said antibody or antigen binding fragment thereof being an isolated monoclonal  
9   antibody or antigen binding fragment thereof which binds to said expressed CD63  
10   antigenic moiety, said antigenic moiety characterized as being bound by an antibody  
11   having the identifying characteristics of a monoclonal antibody encoded by the clone  
12   deposited with the ATCC as PTA-4890;  
  
13          contacting said isolated monoclonal antibody or antigen binding fragment  
14   thereof with said cell sample; and  
  
15          determining binding of said isolated monoclonal antibody or antigen binding  
16   fragment thereof with said cell sample;  
  
17          whereby said cells which express a CD63 antigenic moiety which specifically  
18   binds to an isolated monoclonal antibody encoded by the clone deposited with the  
19   ATCC as PTA-4890, or antigen binding fragment thereof are isolated by said binding  
20   and their presence in said cell sample is confirmed.

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1           Claim 32.     The process of claim 31 wherein the cell sample is obtained  
2     from a tumor originating in a tissue selected from the group consisting of colon,  
3     ovarian, lung, prostate and breast tissue.

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5           Claim 33. A method of extending survival and/or delaying disease progression  
6     by treating a human tumor in a mammal, wherein said tumor expresses an antigen  
7     which specifically binds to a monoclonal antibody or antigen binding fragment thereof  
8     which has the identifying characteristics of a monoclonal antibody encoded by a clone  
9     deposited with the ATCC as accession number PTA-4890 comprising administering to  
10    said mammal said monoclonal antibody in an amount effective to reduce said  
11    mammal's tumor burden, whereby disease progression is delayed and/or survival is  
12    extended.

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15           Claim 34. The method of claim 33 wherein said antibody is conjugated to a  
16    cytotoxic moiety.

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18           Claim 35. The method of claim 33 wherein said cytotoxic moiety is a  
19    radioactive isotope.

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21           Claim 36. The method of claim 33 wherein said antibody activates complement.

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1           Claim 37. The method of claim 33 wherein said antibody mediates antibody  
2 dependent cellular cytotoxicity.

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4           Claim 38. The method of claim 33 wherein said antibody is a murine antibody.

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6           Claim 39. The method of claim 33 wherein said antibody is a humanized  
7 antibody

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9           Claim 40. The method of claim 33 wherein said antibody is a chimerized  
10 antibody.

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